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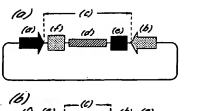
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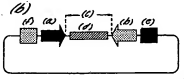
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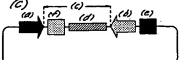
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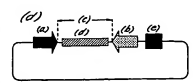


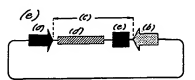


(a): promoter 1
(b): promoter 2



(a): Terminator 1
(f): Terminator 2





(57) Abstract: Vector constructs useful in the expression of double-stranded RNA. The constructs are particularly useful for expression of double-stranded RNA in vitro and in vivo.



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VECTOR CONSTRUCTS

Field of the invention

The invention relates to improved vector constructs for use in the expression of double-stranded RNA, particularly for use in the expression of double-stranded RNA in vitro and in vivo.

Background to the invention

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Since the advent of double-stranded RNA inhibition (RNAi) as a tool for controlling gene expression, as described in WO 99/32619 and WO 00/01846, there has been recognised a need for specialised vectors designed for the production of double-stranded RNA (dsRNA).

Cloning vectors designed to produce high levels of dsRNA have been previously described by Plaetinck et al. (WO 00/01846) and Timmons et al. Nature, 395:854 (1998). These vectors generally contain a multiple cloning site (MCS) into which target DNA fragments can be cloned flanked by two opposable transcriptional promoters. Essentially, these three components (Promoter 1, MCS and Promoter 2) make up the entire system. In the appropriate expression system, the DNA cloned into the MCS may be transcribed in both directions, leading to the production of two complementary RNA strands.

A disadvantage of the known systems is that not only the cloned fragment is transcribed. Read-through of the RNA polymerase will result in transcription of the entire vector, and this also in both directions. As only transcription of the cloned DNA fragment will result in active dsRNA for RNAi purposes, transcription of the vector part results in useless,

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inefficient RNA. More specifically, 80% of these transcripts can be considered as non-specific and thus non-effective.

The large amounts of non-specific RNA generated 5 by the prior art plasmid and expression systems results in some undesirable side effects. First, in RNAi protocols based on introduction of dsRNA into C. elegans via a food organism such as E. coli which expresses the dsRNA (see WO 00/01846), large RNA strands are considered to be toxic for the food 10 organism. As a result, high amounts of RNA accumulating in E. coli cause a significant part of the population to die. Second, and probably more important, is the reduction of inhibition potential. The presence of large amounts of non-specific dsRNA 15 causes a competitive environment for the specified sequences. The potential of the template-specified dsRNA sequences to inhibit the targeted protein expression in, for instance, C. elegans cells is 20 reduced by the presence of these large non-specific regions. Such an inhibition by non-specific dsRNA has also been shown in Drosophila by Tushl et al., Genes & Development 13:3191-3197 (1999). Not only the potential to inhibit gene expression is affected, but also the amount of specific dsRNA produced is limited. Third, transcription of the vector backbone part, more particularly transcription of the origin of replication and related structures, results in plasmid instability and plasmid reorganisation, leading to reduced production of dsRNA. This relatively low concentration of effective dsRNA in turn leads to inefficient RNAi.

To conclude, the previously described vectors have following shortcomings: they are toxic to the

feeding organism, a greater proportion of the transcripts produced are non-specific, the inhibitory potential of the dsRNA is reduced by the presence of non-specific regions, a high incidence of plasmid reorganizations and loss of plasmid from the feeding organism. It is therefore an object of the present invention to provide improved vectors for the production of dsRNA which avoid the disadvantages of the prior art vectors.

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Vectors for use in the in vitro synthesis of RNA transcripts, for example the production of RNA probes, have been known and commonly used in the art for some time (see for example F. M. Ausubel et al. (eds.), Current Protocols in Molecular Biology, John Wiley & Sons, Inc. (1994); Jendrisak et al, Vectors for in vitro production of RNA copies of either strand of a cloned DNA sequence, US 4,766072). In standard in vitro transcription protocols the problem of readthrough transcription of vector sequences is generally avoided by linearizing the transcription vector at restriction site positioned at the 3' end of the desired transcript. However, this solution is not appropriate for in vivo transcription or for the production of dsRNA where it is important that the template is transcribed in both directions.

The inventors now propose a novel solution to the problems encountered with the prior art vectors for the production of dsRNA, based on the use of transcription terminators. Generally the solution consists of the use of at least one transcription terminator operably linked to at least one promoter, wherein the terminator stops the transcription initiated by the promoter. Any DNA fragment inserted between the 3'end of the promoter and the 5' end of

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the terminator will then be transcribed, without the unwanted transcription of the vector backbone. Preferentially the vector consists of two promoters and two terminators, as further described below.

Therefore, in accordance with a first aspect of the invention there is provided a DNA construct comprising two opposable promoters flanking an interpromoter region, the construct further comprising at least one transcription terminator positioned transcriptionally downstream of one of the said promoters In particular, the invention provides for: a DNA construct comprising:

- a) a first promoter and
- b) a second promoter,
- in which the first and second promoter are in opposite orientation to each other and define:
 - c) an inter-promoter region positioned downstream of the 3' end of the first promoter and downstream of the 3' end of the second promoter;
- and which DNA construct further comprises:
 - d)at least one cloning site positioned in the inter-promoter region; and
 - e) a first transcription terminator, positioned (as seen from the 3' end of the first promoter) downstream of the first promoter and downstream of the at least one cloning site, wherein the first transcription terminator is operably linked to the first promoter.

The inter-promoter region can also further be defined as: the DNA region between the 3' end of the first promoter and the 3' end of the second promoter, and which is downstream of the first promoter, and which is downstream of the second promoter, and which preferably does not contains the 5' end of the first

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promoter and of the second promoter. The opposable first promoter and second promoter drive expression directional from their 5' ends to their 3' ends starting transcription downstream of their 3' ends, 5 thus providing transcription of both strands of any nucleotide sequence(s) present in the inter-promoter region.

The two promoters present in the DNA construct of

the invention may be identical or they may be different and may be of essentially any type. The precise nature of the promoters used in the construct may be dependent on the nature of the expression system in which the construct is expected to function (e.g. prokaryotic vs eukaryotic host cell). Bacteriophage promoters, for example the T7, T3 and SP6 promoters, are preferred for use in the constructs of the invention, since they provide advantages of high level transcription which is dependent only on binding of the appropriate RNA polymerase. Each of these promoters can independently by chosen. The phage promoters can also function in a wide variety of host systems, i.e. both prokaryotic and eukaryotic hosts, provided that the cognate polymerase is present in the host cell.

The arrangement of two "opposable" promoters flanking an inter-promoter region such that transcription initiation driven by one of the promoters results in transcription of the sense strand of the inter-promoter region and transcription 30 initiation driven by the other promoter results in transcription of the antisense strand of the interpromoter region is an arrangement well known in the art, for example, in the pGEM7 series of vectors from Promega Corp., Madison WI, USA.

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The DNA constructs of the invention differ from those of the prior art because of the presence of at least one transcription terminator positioned transcriptionally downstream of one of the promoters. The transcription terminator may be uni- or bidirectional, the choice of uni- vs bi-directional terminators being influenced by the positioning of the terminator(s) within or outside the inter-promoter region, as explained below. The terminator may be of prokaryotic, eukaryotic or phage origin. 10 Bacteriophage terminators, for example T7, T3 and SP6 terminators, are particularly preferred. The only requirement is that the terminator must be capable of causing termination of transcription initiating at the promoter relative to which it is transcriptionally 15 downstream. In practice, these means that the promoter and terminator must form a 'functional combination', i.e. the terminator must be functional for the type of RNA polymerase initiating at the promoter. By way of example, a eukaryotic RNA pol II 20 promoter and a eukaryotic RNA pol II terminator would generally form a functional combination. selection of a functional combination is particularly important where bacteriophage promoters and terminators are to be used in the constructs of the 25 invention, since the phage promoters and terminators are both polymerase-specific. To form a functional combination both the promoter and the terminator should be specific for the same polymerase, e.g. T7 promoter and T7 terminator, T3 promoter and T3 terminator etc.

In one embodiment, the DNA construct of the invention may comprise a single transcription terminator, positioned (as seen from the 3' end of the

first promoter) downstream of the first promoter and downstream of the at least one cloning site, wherein the first transcription terminator is operably linked to the first promoter, wherein the single transcription terminator is positioned in the interpromoter region

In an alternative arrangement, the DNA construct comprises a single transcription terminator positioned outside of the inter-promoter region. In a still 10 further embodiment, the DNA construct may comprise two transcription terminators, each one of which is positioned transcriptionally downstream of one of the two promoters. In this arrangement, one or both of the terminators may be positioned within the interpromoter region. These various embodiments of the DNA constructs of the invention will be more fully described below, with reference to the accompanying drawings. The position of a first transcription terminator outside the inter-promoter region may also 20 be further defined as, i.e. such that a first transcription terminator is positioned (as seen from the 3' end of the first promoter) downstream of the first promoter, downstream of the at least one cloning site, and downstream of the 5' end of the second promoter.

The position of a second transcription terminator outside the inter-promoter region may also be further defined as, i.e. such that a second transcription terminator positioned (as seen from the 3' end of the second promoter) downstream of the second promoter, downstream of the at least one cloning site, and downstream of the 5' end of the first promoter.

Moreover, when the terminator is not located in the inter-promoter region, the distance between the 5'

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end of the first promoter and 3' end of the second terminator, or the distance between the 5' end of the second promoter and the 3' end of the first terminator is preferably small, i.e. such that the 3' end of the first transcription terminator is separated from the 5' end of the second promoter by no more than 2000 nucleotides, preferably no more than 1000 nucleotides, more preferably no more than 500 nucleotides, even more preferably no more than 200 nucleotides, especially preferably no more than 100 nucleotides, more especially preferable no more than 50 nucleotides, even more especially preferably no more than 20 nucleotides, particularly preferably no more than 10 nucleotides, more particularly preferably no more than 10 nucleotides, more particularly preferably no more than 6 nucleotides.

Furthermore, when the second transcription terminator is located outside of the inter-promoter region, preferably the 3' end of the second transcription terminator is separated from the 5' end of the first promoter by no more than 2000 nucleotides, preferably no more than 1000 nucleotides, more preferably no more than 500 nucleotides, even more preferably no more than 200 nucleotides, especially preferably no more than 100 nucleotides, more especially preferably no more than 50 nucleotides, even more especially preferably no more than 20 nucleotides, particularly preferably no more than 10 nucleotides, more particularly preferably no more than 10 nucleotides, more particularly preferably no more than 6 nucleotides

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As defined above the term 'inter-promoter region' refers to all of the DNA sequence between the two promoters. As explained above, in certain embodiments of the invention the transcription terminator(s) may be sited within the inter-promoter region. The inter-

promoter region may, advantageously, comprise a sequence of nucleotides forming a template for dsRNA production. The precise length and nature of this sequence is not material to the invention. invention further provides DNA constructs in which the inter-promoter region comprises a cloning site. The function of the cloning site is to facilitate insertion of a DNA fragment forming a template for dsRNA production between the two promoters. Thus, the invention provides a series of cloning vectors which are of general use in the construction of template vectors for dsRNA production. Also encompassed within the scope of the invention are vectors derived from the cloning vectors which have a DNA fragment inserted into the cloning site. 15

The cloning site may further comprise one or more of the following:

- at least one restriction site, (as known in the art), or one or more further restriction sites,
 e.g. to provide a multiple cloning site(as known in the art),
- a stuffer DNA, e.g., flanked by at least two restriction site, such as two BstXI restriction sites, or two XcmI restriction sites,
- 25 attR1 and attR2 recombination sites,
 - a ccdB nucleotide sequence,
 - a ccdB nucleotide further comprising at least one unique blunt-end restriction site, such as a SrfI restriction site, and/or
- a DNA fragment inserted in the at least one cloning site.All of the DNA constructs provided by the invention may, advantageously, form part of a replicable cloning vector, such as, for example, a plasmid vector. In addition to the opposable

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promoters, inter-promoter region and transcription terminator(s), the vector 'backbone' may further contain one or more of the general features commonly found in replicable vectors, for example an origin of replication to allow autonomous replication within a host cell and a selective marker, such as an antibiotic resistance gene. The selective marker gene (e.g. the antibiotic resistance gene) may itself contain a promoter and a transcription terminator and it is to be understood that these are completely independent of the promoter and terminator elements required by the invention and are not to be taken into consideration in determining whether a particular vector falls within the scope of the invention.

DNA constructs according to the invention may be easily be constructed from the component sequence elements using standard recombinant techniques well known in the art and described, for example, in F. M. Ausubel et al. (eds.), Current Protocols in Molecular Biology, John Wiley & Sons; Inc. (1994), as will be appreciated by one skilled in the art from the following detailed description of the invention and the accompanying Examples.

There follows a detailed description of DNA constructs according to the invention, with reference to the following schematic drawings in which:

Figures 1(a) to 1(e) are schematic representations of several different embodiments of the DNA construct according to the invention illustrating the relative positioning of the promoter and transcription terminator elements.

Figure 2(a) is a schematic representation of a prior

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art vector included for comparison purposes.

Figures 2(b) to 2(e) are schematic representations of several further embodiments of the DNA construct according to the invention illustrating the use of different cloning sites in the inter-promoter region.

Referring to the Drawings, Figure 1(a) schematically illustrates a first DNA construct according to the invention which is a plasmid vector comprising two opposable promoters; a first promoter a) and and second promoter b) 2 flanking an interpromoter region c) , which inter-promoter region is downstream of the 3' of the first promoter, and down stream of the 3' end of the second promoter. The first promoter and the second promoter may be identical or different. This embodiment comprises a first transcription terminators e) and a second transcription terminator f)both of which are positioned within the inter-promoter region. embodiment, the first terminator and the second terminator are preferentially uni-directional terminators.

A DNA fragment may be inserted in the at least one cloning site d). Such fragment is subject to transcription directed by the first promoter a) and the second promoter b) (i.e. transcription of both strands), resulting in the generation of two RNA fragments which may combine to double-stranded RNA of the inserted DNA fragment (both in vitro and in vivo).

Any desired DNA sequence, such as a genomic DNA sequence, or a cDNA sequence or any other coding sequence, may be inserted in the at least one cloning site. Without being limited to any specific

explanation, it is assumed that when a) and e) form a functional combination, RNA polymerase which initiates transcription at a) will transcribe the inter-promoter region including the at least cloning site and the DNA fragment inserted in the at least cloning site and will be terminated when it reaches e). Similarly, RNA polymerase which initiates transcription at b) will transcribe the inter-promoter region including the at least one cloning site and the DNA fragment inserted 10 in the at least one cloning site and will terminate when it reaches f). The terminators cause the RNA polymerase to pause, stop transcription and fall off the template. This prevents the unlimited transcription of the vector backbone, and reduces the unspecific transcription of non-essential DNA.

The inter-promoter region further comprises a sequence of nucleotides corresponding to a target for double-stranded RNA inhibition. This sequence is designated 'TF'' for target fragment. It is this sequence which, when transcribed into dsRNA, will be responsible for specific double-stranded RNA inhibition of a target gene. The target fragment may be formed from a fragment of genomic DNA or cDNA from the target gene. Its precise length and nucleotide sequence are not material to the invention.

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In the arrangement shown in Figure 1(a) the two terminators are positioned on either side of the TF within the inter-promoter region. Each of the terminators is positioned transcriptionally downstream of one of the promoters, the fist terminator e) is transcriptionally downstream of first promoter a) and the second terminator f) is transcriptionally downstream of the second promoter b). Assuming that a) and e) form a functional combination, as described

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above, RNA polymerase which initiates transcription at a) will transcribe the inter-promoter region up to and including TF and will be terminated when it reaches e). Similarly, RNA polymerase which initiates transcription at b) will transcribe the inter-promoter region up to and including TF on the opposite strand and will terminate when it reaches f). The terminators cause the RNA polymerase to pause, stop transcription and fall off the template. This prevents the unlimited transcription of the vector backbone, and reduces the unspecific transcription of non-essential DNA.

The transcripts generated from this vector may, depending on the precise placement of the terminators in the vector, be almost completely specific dsRNAs 15 corresponding to the TF region. Through the direct placement of the terminator sequences at the downstream end of the TF region on both sides of the inserted DNA fragment, the amount of material transcribed is completely reduced to the 20 template-specified sequences. Therefore, a higher amount of specific dsRNA is obtained. Furthermore the constructs are now also more stable, due to the non-transcription of the vector backbone. The latter 25 characteristic (stability), combined with the now relatively higher specific transcription rate, provides a system adapted to synthesise higher amounts of specific short dsRNA strands. This proportionally higher amount of transcript, resulting in high concentrations of specific dsRNA, enhances the 30 inhibitory effect in RNAi protocols. In RNAi protocols based on expression of dsRNA in a food organism, toxicity for the feeding organisms due to high RNA expression is brought to a minimal level by

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use of this vector.

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A specific example of a vector of the type illustrated in Figure 1(a), considered by the inventors to be the optimal arrangement for RNAi applications, is plasmid pGN9 described in the accompanying Examples. The transcriptional terminators used in pGN9 are T7 RNA polymerase specific terminators, since the vector contains two opposable T7 promoters. However, other systems could be used such as an expression system based on the T3 or SP6 promoter, terminator and polymerase or other prokaryotic or eukaryotic promoters and terminators.

Figure 1(b) illustrates schematically a further 15 DNA construct according to the invention which is a plasmid vector comprising two opposable promoters a) and b) flanking an inter-promoter region c). vector also comprises two transcription terminators e) and f) but in this arrangement the two terminators are positioned outside of the inter-promoter region, in fact the terminator elements now flank the two promoters. The arrangement is such that e) is transcriptionally downstream of a) whilst f) is transcriptionally downstream of b). Here again e) terminates the transcription initiated by a), whilst f) terminates the transcription initiated by promoter Placement of the terminators outside of d)allows the use of bi-directional terminators as well as unidirectional terminators, in contrast to the arrangement in Figure 1(a) where uni-directional terminators are preferred because of the placement of the terminators between the promoters. A number of bi-directional terminators which could be used in accordance with the invention are known in the art.

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Generally these are observed to be polymerase nonspecific.

The embodiment shown in Figure 1(b) provides essentially the same advantages as that shown in

5 Figure 1(a) over the prior art vectors for dsRNA production. The vector shown in Figure 1(b) will lead to the production of transcripts which are slightly longer, including the promoter regions. This relatively small difference in the length of the

10 transcript and hence the formed dsRNA will not severely affect the efficacy in an RNAi system.

The position of the terminators and promoter in the example as shown in figure 1 (b) are preferably placed at close proximity, such that the 5' end of the promoters are separated from the 3' end of the transcription terminators by no more than 2000 nucleotides, preferably no more than 1000 nucleotides, more preferably no more than 500 nucleotides, even more preferably no more than 200 nucleotides, even more preferably no more than 100 nucleotides, more especially preferably no more than 50 nucleotides, even more especially preferably no more than 20 nucleotides, particularly preferably no more than 10 nucleotides, more particularly preferably no more than 10 nucleotides, more particularly preferably no more than 6 nucleotides.

Figure 1(c) illustrates schematically a further DNA construct according to the invention which is a plasmid vector comprising two opposable promoters a) and b) flanking an inter-promoter region c). In this embodiment one terminator (in this case e)) is positioned within the c) and the other (f)) is positioned outside c). Again, e) terminates transcription initiated by a) and f) terminates

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transcription initiated by b). This arrangement may provide a useful solution to the problem of one of the terminators interfering with polymerase activity in the other direction (e.g. f) interferes with b) initiated transcription). This vector essentially provides the same advantages as the vector variations shown in Figure 1(a) and Figure 1(b) over the prior art. The relatively small difference in the length of the transcript due to the transcription of one of the promoters will not significantly affect the efficacy in RNAi systems. This more particularly the case when the terminator that is located outside of the interpromoter region c) is at close proximity of the promoter, as defined above.

15 Figures 1(d) and 1(e) illustrate schematically two further DNA constructs according to the invention which are both plasmid vectors comprising two opposable promoters a) and b) flanking an interpromoter region c). These embodiments comprise a single terminator only. In the arrangement shown in Figure 1(d) a single terminator e) which terminates transcription from a) is placed outside of c). The placement of the terminator outside of the IPR allows the use of both a bi-directional terminator or a unidirectional terminator in this system. 25 embodiment shown in Figure 1(d) e) is placed within the c). a) should therefore preferably be a unidirectional terminator.

Further embodiments of the DNA construct according to the invention are illustrated schematically in Figures 2(b) to 2(e).

These embodiments are all plasmid cloning vectors, based upon the optimal arrangement of promoters and terminators shown in Figure 1(a), and

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described above, containing cloning sites to facilitate the insertion of a DNA fragment into the at least on cloning site.

These embodiments are all plasmid cloning vectors, based upon the optimal arrangement of promoters and terminators shown in Figure 1(a), containing cloning sites to facilitate the insertion of a target DNA fragment into the inter-promoter region.

10 Figure 2(a), which is a schematic representation of a prior art cloning vector, is included for comparison purposes. This vector comprises two opposable promoters a) and b), which may be identical or different, flanking a multi-cloning site (MCS).

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Figure 2(b) illustrates a first type of plasmid cloning vector according to the invention. The vector contains a first opposable promoters a) and a second opposable promoter b) flanking an inter-promoter region. The inter-promoter region can further be defined as: the DNA region between the 3' end of the first promoter and the 3' end of the second promoter, and which is downstream of the first promoter, and which is downstream of the second promoter, and which preferably does not contains the 5' end of the first promoter and of the second promoter. The interpromoter promoter region further comprises terminators e) and f) flanking a multi-cloning site The MCS comprises at least one individual restriction sites, an preferably more than one restriction site as known in the art, any of which may be used for insertion of a DNA fragment.

Figure 2(c) illustrates a further type of plasmid cloning vector according to the invention. This vector again contains opposable promoters a) and b)

flanking an inter-promoter region comprising terminators e) and f). In this embodiment, a) and b) flank a cloning site which is adapted for facilitated cloning of PCR fragments, comprising a stuffer DNA flanked by two identical restriction sites, in this case BstXI sites. The specific sequence of the stuffer DNA is not essential, provided that said stuffer DNA does not interfere with the desired effect and/or the desired activity of the DNA constructs of the invention. A specific example of a vector according to this aspect of the invention described herein is plasmid pGN29.

The cloning of PCR products using BstXI recognition sites and BstXI adaptors is generally known in the art. The BstX1 adaptors are commercially obtained, such as from Invitrogen (Groningen, the Netherlands). These adaptors are non-palindromic adapters designed for easier and efficient cloning of PCR products into vectors. These use of these adaptors reduces the concatemerization of adapters or insert DNA by having non-complementary (CACA) overhangs. The stuffer DNA is included merely to allow easy differentiation between BstXI cut and uncut vector on the basis of size. Its precise length and sequence are not of importance.

Figure 2(d) illustrates a further type of plasmid cloning vector according to the invention. This vector again contains opposable promoters a) and b) flanking an inter-promoter region comprising terminators e) and f). In this embodiment, a) and b) flank a cloning site which facilitates "High Throughput" cloning based on homologous recombination rather than restriction enzyme digestion and ligation. As shown schematically in Figure 2(d), the cloning

site comprises attR1 and attR2 recombination sites from bacteriophage lambda flanking a gene which is lethal to E. coli, in this case the ccdB gene.

An alternative cloning method of DNA fragments into this vector, (not shown in Figure 2 (d)), consists of a variant of this vector, wherein the *ccd*B DNA sequence further comprises at least one unique restriction site, preferably the at least unique restriction site is a blunt-end restriction site, such as a *SrfI* restriction site. Insertion of a DNA fragment in the at least unique restriction, results in inactivation of the ccdB gene, and hence in inactivation of the lethal ccdB gene.

A further variant of a vector a shown in Figure 2(d) in which the attR1 and the attE2 are not present. Such a vector comprises an at least one cloning site, said at least one cloning site consisting of a ccdB sequence, said ccdB sequence further comprising at least one unique restriction site, preferably the at least unique restriction site is a blunt-end restriction site, such as a SrfI restriction site. Insertion of a DNA fragment in the at least unique restriction, results in inactivation of the ccdB gene, and hence in inactivation of the lethal ccdB gene.

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These cloning sites comprising the ccdB nuclotide sequence and/or the attR sites (R1 and/or R2) are derived from the Gateway™ cloning system commercially available from Life Technologies, Inc. The Gateway™ cloning system has been extensively described by Hartley et al. in WO 96/40724 (PCT/US96/10082). A specific example of a vector according to this aspect of the invention described herein is pGN39.

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Figure 2(e) and 2(f) illustrate a still further type of plasmid cloning vector according to the invention. This vector again contains opposable promoters a) and b) flanking an inter-promoter region c) comprising terminators e) and f). In the embodiment shown in Figure 2(e), e) and f) flank a cloning site which facilitates "high throughput" cloning of PCR products by TATM cloning. This cloning site comprises a stuffer DNA flanked by two identical 10 restriction sites for an enzyme which generates overhanging T nucleotides. In this case the restriction sites are XcmI sites, but other sites which are cleaved to generate overhanging T nucleotides could be used with equivalent effect. 15 overhanging T nucleotides facilitate cloning of PCR products which have a overhanging A nucleotide. This principle is known as TA™ cloning. The cut vector with overhanging T nucleotides can be "topomerized" to generate a cloning vector of the type shown 20 schematically in Figure 2(f), by linking the enzyme topoisomerase to the overhanging T nucleotides. The resulting vector also facilitates the cloning of PCR products by the principle known as $TOPO^{TM}$ cloning.

Both the TOPO[™] and TA[™]cloning systems, although not for the vectors described in this invention, are commercially available from Invitrogen. The TOPO[™] cloning system has extensively been described by Shuman in WO 96/19497 (PCT/US95/16099). The TA[™] cloning system has extensively been described by Hernstadt et al. in WO 92/06189 (PCT/US91/07147).

It will be readily appreciated by the skilled reader that whilst Figures 2(b)-2(f) illustrate the inclusion of different cloning sites into a vector of the type illustrated in Figure 1(a), these cloning

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sites could be included into any of the DNA constructs of the invention, including those illustrated schematically in Figures 1(b) to 1(e)

5 Application of the DNA constructs of the invention in RNAi technology.

As aforementioned, a major application of the DNA constructs/vectors of the invention is in the production of double stranded RNA for use in RNAi technology. In particular, the constructs are useful in *in vivo* RNAi protocols in the nematode worm *C*. elegans.

In C. elegans, RNAi has traditionally been performed by injection dsRNA into the worm. Fire et al. describes these methods extensively in International Application No. WO 99/32619. both strands of RNA are produced in vitro using commercially available in vitro transcription kits. Both strands of RNA are allowed to form dsRNA, after 20 which the dsRNA is injected into C. elegans. The new vector system developed by the present inventors is a drastic improvement on this traditional method. First, the RNA can be produced in one step, for instance by using two identical promoters such as 25 in the vector pGN9. Second, and more important, due to the presence of terminators the transcripts, and hence the formed dsRNA, will be more specific as only the cloned target fragment will be transcribed. This will result in a more efficient RNAi.

A further method to perform RNAi experiments in C. elegans has been described by Plaetinck et al. in WO 00/01846. In this method C. elegans worms are fed with bacteria which produce dsRNA. The dsRNA passes the gut barrier of the worm and induces the same RNAi

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as if the dsRNA has been injected. For these experiments, the preferred *E. coli* strain is HT115 (DE3), and the preferred *C. elegans* strain is nuc-1;gun-1. The improved vectors provided by the invention also improve the efficiency of RNAi in this method, as shown in the example below, as only effective dsRNA is produced.

Another method to perform RNAi has also been described by Plaetinck et al. in WO 00/01846. In short, this method is based on the production of dsRNA in the worm itself. This can be done by using worm promoters in the described vectors, or by using a transgenic worm expressing a polymerase specific for non-C. elegans promoters present in the vector, such that this polymerase drives transcription of the dsRNA. The promoters will preferentially be selected from the known bacteriophage RNA promoters, such as T7 or T3 or SP6 RNA promoters, which provide the advantage of a high level of transcription dependent only on the binding of the cognate polymerase.

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Plasmid vector DNA can be introduced into the worm by several methods. First, the DNA can be introduced by the traditional injection method (Methods in Cell Biology, Vol 48, C. elegans Modern Biological Analysis of an organism, ed. by Epstein and Shakes). Second, the DNA can be introduced by DNA feeding. As has been shown by Plaetinck et al. in WO 00/01846, plasmid DNA can be introduced into the worm by feeding the worm with an *E. coli* strain that harbors the plasmid. Preferentially the *E. coli* strain is OP50, or MC1061 or HT115 (DE3) but any other strain would suit for this purpose. The *C. elegans* strain is preferentially a nuc-1 mutant strain or a nuc-1; gun-1 strain. The plasmid DNA from the *E. coli*

passes the gut barrier and is introduced into the nematode, resulting in the expression of dsRNA. As with the other RNAi methods described above, the use of the new vector system will enhance the RNAi by the production of only specific dsRNA.

The invention will be further understood with reference to the following experimental Examples, together with the following additional Figures in which:

- Figure 3 is a representation (plasmid map) of pGN1.
- Figure 4 is a representation (plasmid map) of pGN9.

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- Figure 5 illustrates the nucleotide sequence of a fragment of plasmid pGN1, annotated to show the positions of the opposable T7 promoters.
- 20 Figure 6 depicts the nucleotide sequence of the T7 transcription terminator.
- Figure 7 illustrates the sequences of oligonucleotides oGN27, oGN28, oGN29 and oGN30 used to insert T7 transcription terminators into pGN1. The positions of the T7 pol terminator sequences and of various restriction sites are marked.
- 30 Figure 8 illustrates the nucleotide sequence of a fragment of plasmid pGN9, annotated to show the positions of the opposable T7 promoters and the T7 transcription terminators.

Figure 9 (a) is a representation (plasmid map) of pGN29; (b) is a representation (plasmid map) of pGN39; (c) is a representation (plasmid map) of the plasmid TopoRNAi.

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- Figure 10 shows the complete nucleotide sequence of plasmid pGN9.
- Figure 11 shows the complete nucleotide sequence of plasmid pGN29.
 - Figure 12 shows the complete nucleotide sequence of plasmid pGN39.
- 15 Figure 13 shows the complete nucleotide sequence of plasmid TopoRNAi.
 - Figure 14 shows the complete sequence of plasmid pGN49A.

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- Figure 15 shows the complete sequence of plasmid pGN59A.
- Figure 16 is a representation (plasmid map) of pGN49A.

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Figure 17 is a representation (plasmid map) of pGN59A.

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Example 1-Vector construction.

The starting point for construction of the vectors exemplified herein was plasmid pGN1. This plasmid, described in the applicant's co-pending International Application No. WO 00/01846, contains two opposable T7 promoters flanking a multi-cloning site.

Vector construction was carried out according to standard molecular biology techniques known in the art and described, for example, in F. M. Ausubel et al. (eds.), Current Protocols in Molecular Biology, John Wiley & Sons, Inc. (1994).

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1) Construction of pGN9

pGN1 was first digested with restriction enzymes EcoRI and KpnI. Oligonucleotides oGN27 and oGN28 (Figure 7) were annealed to generate a double stranded fragment which was then ligated into the EcoRI/KpnI cut vector. The resulting plasmid was re-digested with XbaI and HindIII. Oligonucleotides oGN29 and oGN30 were annealed to generate a double-stranded fragment which was then annealed into the XbaI/HindIII cut vector. The resulting vector was designated pGN9 (Figures 4 and 10).

pGN29 (Figure 9(a); Figure 11) was generated by replacing the MCS in pGN9 with a stuffer DNA flanked by BstXI sites. BstXI adapters are commercially

2) Construction of further cloning vectors

by BstXI sites. BstXI adapters are commercially available from Invitrogen (Groningen, the Netherlands).

pGN39 (Figure 9 (b); Figure 12) generated by following steps; pGN29 was digested with BstXI. BstXI adapters (Invitrogen, Groningen, The Netherlands) were ligated to Cassette A provided by the GATEWAYTM system (Life Technologies, Inc.). Cassette A contains attR1, CmR, CcdA, CcdB, attR2. The Cassette A with the adapters where then ligated into the digested pGN29, resulting in pGN39A. pGN39A contains a unique SrfI site in the ccdB gene.

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The TopoRNAi vector (figure 9 (c); figure 13) was generated in the following way; pGN29 was digested with BstXI. Using PCR with the primers oGN103 and oGN104 and template pCDM8 (Invitrogen, Groningen, The Netherlands), a stuffer was generated which includes XcmI sites. Onto the PCR product, BstXI adapters were ligated, and the resulting ligation product was ligated in the BstXI digested pGN29 vector resulting in the TopoRNAi vector.

ogN103:5'TACCAAGGCTAGCATGGTTTATCACTGATAAGTTGG 3' ogN104:5'TACCAAGGCTAGCATGGGCCTGCCTGAAGGCTGC 3'

PGN49A was constructed to insert an additional unique non-blunt restriction site and to delete the CmR gene pGN39. Overlap PCR was used. A first PCR was performed with primers oGN126 and oGN127 and PGN39A as template. Using primers oGN128 an oGN129 and the same template a second fragment was generated. Overlap PCR using the resulting fragments and primers oGN126P and oGN129P resulted in a final PCR product. To this final PCR

- 27 -

product, BstXI adapters were ligated, and the ligation product was ligated into pGN29 digested with BstXI. The resulting vector was designated pGN49A.

of the pGN49A cloning vector, such vector should contain the T7 promoters, but not the T7 terminators. For this, the XbaI insert of pGN49A was isolated and cloned in pGN1 digested with the same restriction enzyme. The resulting vector was designated pGN59A.

ogN126 pGATCTGGATCCGGCTTACTAAAAGCCAGATAACAGTATGC
oGN127 GGAGACTTTATCGCTTAAGAGACGTGCACTGGCCAGGGGGATCACC
oGN128:

5 CCAGTGCACGTCTCTTAAGCGATAAAGTCTCCCGTGAACTTTACCCGGTGG
OGN129 pGCTGTGTATAAGGGAGCCTGACATTTATATTCCCCAG

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Example 2-To illustrate the usefulness of the improved vectors in RNA.

This experiment was designed to illustrate the improved efficiency of the improved vectors of this invention in double-stranded RNA inhibition, as compared to the vectors known from the prior art. A significant increase on the efficacy of the system could be expected, as more effective dsRNA was produced and hence RNAi performed better. The experimental system for this proof of concept experiment was to measure *C. elegans* rescue at 25°C in nuc-1 / pha-1(e2123)ts *C. elegans* mutants by RNAi of sup35 using dsRNA feeding of pGN-2 (-terminator) and pGN-12 (+ terminator), with PGN-1 (empty vector) as a control and dilutor. The pha-1 ts / sup-35 mutation has extensively been described by Schnabel in WO

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99/49066.

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The *nuc-1* mutation in C. elegans provides for a C. elegans strain exhibiting better uptake abilities, such as the uptake of the dsRNA complexes than wild type C. elegans. This mutant is deleted in the major DNAse enzymes, and inventors have proven in earlier co-pending applications that this C. elegans strain results in enhanced RNAi by feeding the nematode with dsRNAs.

The pha-1(e2123)ts mutation provides a mutant *C*. elegans strain with a phenotype of survival at 15°C and lethality at 25°C. This lethality is suppressible by the inhibition of sup-35 expression. RNAi of sup-35 should thus facilitate the rescue of pha-1(e2123)ts at 25°C. The vectors of the present invention, when expressing dsRNA from sup-35, should increase the efficacy of sup-35 RNAi in rescuing pha-1(e2123)ts mutants at 25°C, compared to vectors that do not contain the terminators.

Vector pGN1 was used as empty vector. Vector pGN2 (-terminator) is a vector harboring sup-35 DNA and expressing sup-35 dsRNA when introduced in the appropriate host, the vector has no transcriptional terminators inserted. Vector pGN12 (+ terminator) is a vector as described above, containing the transcriptional terminators, and hence resulting in improved dsRNA production when introduce into an appropriate host. Thus, this vector has two unidirectional transcriptional terminators, both placed inside of the inter-promoter region, and flanking the sup-35 fragment. Use of the latter

vector was expected to increase the efficacy of the system, here meaning a better rescue (survival) of pha-1(e2123)ts mutants at 25°C.

5 Experimental conditions

12-well micro-titer plates were filled with approximately 2 ml of NGM agar per well.

(1 liter of NGM agar: 15g Agar, 1g peptone, 3g NaCl, lml cholesterol solution (5 mg/ml in EtOH), with sterile addition after autoclaving of 9.5 ml 0.1M CaCl₂, 9.5 ml 0.1 ml MgSO₄, 25 ml 1M KH₂PO₄/K₂HPO₄ buffer pH 6, Ampicillin (100 μg/l), 5ml 0,1M IPTG and 5 ml nystatin solution (dissolved 10 mg/ml in 1:1 EtOH:CH₃COONH₄ 7.5 M)

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The dried plates were spotted with approximately 50 µl of an overnight culture of bacteria HT115 (DE3) (Fire A, Carnegie Institution, Baltimore, MD) transformed with the plasmids. Individual nematodes at the L4 growth stage were then placed in single wells at day 1. In each well 1 nematode (P1). At day two, the P1 nematodes were placed to a new well and left to incubate for a day. The same procedure was repeated at day 3. All plates were further incubated at 25°C to allow offspring to be formed. Sup35 RNAi induced survival (rescue) was measured by counting the offspring.

Results

RNAi experiment in C. elegans nuc-1/pha-1(e2123)ts mutants by feeding with E. coli expressing sup-35 dsRNA.

Set up:

pGN1 as control

pGN2 (sup 35 - Term.)

pGN12 (sup 35 + Term.)

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pGN2 + pGN1 dilutions 1/2, 1/4, 1/8, 1/16, 1/32 pGN12 + pGN1 dilutions 1/2, 1/4, 1/8, 1/16, 1/32

10 Conditions:

Incubation temperature 25°C

Readout:

Count offspring (adult hermaphrodites)

pGN1 (control)

Day 1	0	0	0	0
Day 2	0	0	0	. 0
Day 3	0	0	0	0

pGN2 (undiluted)

Day 1	12	4	48	32
Day 2	24	23	80	85
Day 3	5.	0	9	16

pGN 2+1, 1/2 dilution

Day 1	0	7	0	2
Day 2	9	10	0	3
Day 3	0	2	0	0

pGN12 (undiluted)

Day 1	16	29	37	14
Day 2	27	22	57	2
Day 3	1	2	4	1

pGN 12+1, 1/2 dilution

Day 1	22	28	103	61
Day 2	3.6	45	53	40
Day 3	3	. 3	25	1

pGN 2+1, 1/4 dilution

Day 1	28	23	0	0
Day 2	6	3	0	0
Day 3	0	0	0	0

pGN 12+1, 1/4 dilution

Day 1	*	6	36	5
Day 2		24	55	3
Day 3			•	

pGN 2+1, 1/8 dilution

Day 1	0	0	4	0
Day 2	0	O ₀	11	0
Day 3	0	0	0	0

pGN 12+1, 1/8 dilution

Day 1	31	12	16	38
Day 2	4	5	37	4
Day 3	0	0	2	1

pGN 2+1, 1/16 dilution

Day 1	0	0	0	0
Day 2	0	0	0	1 little
Day 3	0	0.	0	0

pGN 12+1, 1/16 dilution

Day 1	1	0	0 .	0
Day 2	2	0	0	1
Day 3	0	1	1	1

pGN 2+1, 1/32 dilution

		•		
Day 1	0	0	0	0
Day 2	0	0	0	0
Day 3	0	0	0	0

pGN 12+1, 1/32 dilution

Day 1	0	0	1	0
Day 2	0	L2	3	0
Day 3	2	0	L3- L4	0

* mother died

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Conclusions

generation of RNAi.

WO 01/88121

As expected, worms fed by bacteria harboring pGN1, did not result in the viable offspring, due to the lethal effect of the pha-1 mutation at this temperature. 5 Feeding the nematodes with E. coli harboring pGN2 or pGN12 both result in viable offspring. This is due to the feeding of the worm with dsRNA from sup-35. remarkable difference between the two feeding experiments can be seen in the dilution series. When 10 diluting the bacteria harboring pGN2 with bacteria harboring pGN1, the number of offspring diminishes drastically, even at a low dilution of one to two. This dilution series indicates that high levels of dsRNA are needed to have a proper RNAi induction. In the feeding experiment with bacteria harboring pGN12, 15 significant offspring is still observed at a dilution of one to eight. This indicates that in the bacteria harboring pGN12, much more effective dsRNA is formed. This experiment clearly shows that the addition of 20 terminator sequences in vectors to express dsRNA as described above provide a significant advantage in the

Example 3: Comparison of RNAi efficiency of vectors with and without T7 terminators(pGN49 vs pGN59)

Three different genes have been cloned in the vectors pGN49A and pGN59A. The cloning was performed by amplifying the gene fragments with PfuI DNA polymerase producing blunt ends, facilitating cloning in these vectors. These PCR fragments were cloned into the vectors digested with SrfI. Correct fragment insertion of the clones was checked by PCR. The fragments are chosen such that ds expression and RNAi results in a

lethal phenotype of the offspring. This procedure allows to compare fast and easy the efficiency of the two vectors pGN49 and pGN59 in RNAi.

plasmid	Gene	(acedb)	Vector	backbone5
pGW5	B0511	. 8	pGN49A	
pGW9	C01G8	.7	pGN49A	
pGW11	C47B2	.3	pGN49A	·
pGW17	B0511	.8	pGN59A	
pGW21	C01G8	.7	pGN59A	(
рG ў 23	C47B2	.3	pGN59A	
<u> </u>			\ _	

All the plasmids (pGW-series) are transformed in *E.coli* AB301-105 (DE3) bacteria by standard methodology. The bacteria are then grown in LB/amp at 37°C for 14-18h.

These cultures were centrifuged and the bacterial pellet dissolved in S-complete buffer containing 1mM IPTG and 100 μ g/ μ l ampiciline.

In 96 well plates containing 100 µl S-complete

(containing 1 mM IPTG and 100 µg/µl ampiciline final concentration) and 10 µl of bacteria solution, 3 nematodes were added at each well, the nematodes were at the L1 growth stage.

The plates were incubated at 25°C for 5-6 days. Each

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day the plates are inspected for development of the larvae and the production of F1 offspring.

5 Results

The RNAi was performed in eight-fold for each constructed plasmid. The results show that when T7 terminators are inserted into the vector backbone, the expected phenotype gives a 100% occurrence. When T7 terminators are not used the reproducibility can decrease up to 50%. As in the previous experiment, the results show that the addition of terminators significantly enhances RNAi performance.

	DNA						
	fragment	B0511.8	B0511.8	C01G8.7	C01G8.7	C47B2.3	C47B2.3
	Vector	pGN49A	pGN59A	PGN49A	pGN59A	pGN49A	pGN59A
	Resulting					•	
	plasmid	PGW5	PGW17	PGW9	PGW21	PGW11	PGW23
•	Percentage			•			
	lethal	100	75	100	87.5	100	50
	Percentage						
	offspring	0	25	0	12.5	0	50

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PCR fragment generated by the primers oGN103 and oGN104 on template pCDM8

TACCAAGGCT AGCATGGTTT ATCACTGATA AGTTGG ATAAGTTGGT GGACATATTA TGTTTATCAG TGATAAAGTG TCAAGCATGA CAAAGTTGCA GCCGAATACA GTGATCCGTG CCGGCCCTGG ACTGTTGAAC GAGGTCGGCG TAGACGGTCT GACGACACGC AAACTGGCGG AACGGTTGGG GGTGCAGCAG CCGGCGCTTT ACTGGCACTT CAGGAACAAG CGGGCGCTGC TCGACGCACT GGCCGAAGCC ATGCTGGCGG AGAATCATAC GCTTCGGTGC CGAGAGCCGA CGACGACTGG CGCTCATTTC TGATCGGGAA TCCCGCAGCT 10 TCAGGCAGGC CCATGCTAGC CTTGGTACCA GCACAATGG

Overlap PCR Fragment, which was used to generate 15 pGN49A

gatctggatccggcttactaaaagccagataacagtatgcgtatttgcgcgctg atttttgcggtataagaatatatactgatatgtatacccgaagtatgtcaaaaa gaggtgtgctatgaagcagcgtattacagtgacagttgacagcgacagctatca 20 gttgctcaaggcatatatgatgtcaatatctccggtctggtaagcacaaccatg cagaatgaagcccgtcgtctgcgtgccgaacgctggaaagcggaaaatcaggaa gggatggctgaggtcgcccggtttattgaaatgaacggctcttttgctgacgag ttatcgtctgtttgtggatgtacagagtgatattattgacacgcccgggcga $\verb|cgg| atggtgatcccctggccagtgcacgtctcttaagcgataaagtctccc|$ gtgaactttacccggtggtgcatatcggggatgaaagctggcgcatgatgac caccgatatggccagtgtgccggtctccgttatcggggaagaagtggctgat ctcagccaccgcgaaaatgacatcaaaaacgccattaacctgatgttctqqq gaatataaatgtcaggctcccttatacacagc

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Claims:

- 1. A DNA construct comprising:
- a) a first promoter and
- 5 b) a second promoter, in which the first and second promoter are in opposite orientation to each other and define:
 - c) an inter-promoter region positioned downstream of the 3' end of the first promoter and downstream of the 3' end of the second promoter; and which DNA construct further comprises:
 - d) at least one cloning site positioned in the interpromoter region; and
- e) a first transcription terminator, positioned (as

 seen from the 3' end of the first promoter)

 downstream of the first promoter and downstream of

 the at least one cloning site, wherein the first

 transcription terminator is operably linked to the

 first promoter.

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- 2. A DNA construct according to claim 1, further comprising:
- f) a second transcription terminator positioned (as seen from the 3' end of the second promoter)
- downstream of the second promoter and downstream of the at least one cloning site.

 wherein the second transcription terminator is operably linked to the second promoter.
- 30 3. A DNA construct according to claim 1 or 2, in which the first transcription terminator is positioned in the inter-promoter region.

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4. A DNA construct according to claim 1 or 2, in which the first transcription terminator is positioned (as seen from the 3' end of the first promoter) downstream of the first promoter, downstream of the at least one cloning site, and downstream of the 5' end of the second promoter.

- A DNA construct according to any one of claims 2,
 3 or 4, in which the second transcription terminator is positioned in the inter-promoter region.
- 6. A DNA construct according to any of claims 2, 3
 or 4 in which the second transcription terminator is positioned (as seen from the 3' end of the second promoter) downstream of the second promoter, downstream of the at least one cloning site, and downstream of the 5' end of the first promoter.
- 7. A DNA construct according to any one of claims 4, 5 or 6, in which the 3' end of the first transcription terminator is separated from the 5' end of the second promoter by no more than 2000 nucleotides, preferably no more than 1000 nucleotides, more preferably no more than 500 nucleotides, even more preferably no more than 200 nucleotides, especially preferably no more than 100 nucleotides, more especially preferably no more than 50 nucleotides, even more especially preferably no more than 20 nucleotides, particularly preferably no more than 10

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nucleotides, more particularly preferably no more than 6 nucleotides.

- 8. A DNA construct according to any one of claims 6 or 7, in which the 3' end of the second 5 transcription terminator is separated from the 5' end of the first promoter by no more than 2000 nucleotides, preferably no more than 1000 nucleotides, more preferably no more than 500 nucleotides, even more preferably no more than 10 200 nucleotides, especially preferably no more than 100 nucleotides, more especially preferably no more than 50 nucleotides, even more especially preferably no more than 20 nucleotides, particularly preferably no more than 10 15 nucleotides, more particularly preferably no more than 6 nucleotides.
- A construct according to any one of the preceding
 claims wherein the first and the second promoter are identical.
- 10. A DNA construct according to any one of the claims 1 to 7 wherein the first and the second promoter are non-identical.
- 11. A DNA construct according to claims 8 or 9
 wherein the first promoter and the second
 promoter are independently chosen from T7, T3 or
 SP6 promoters.
 - 12. A construct according to any one of the preceding claims wherein the cloning site comprises at

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least one restriction site.

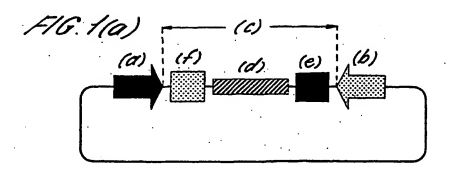
- 13. A DNA according to claim 11 wherein the cloning site comprises at least two restriction sites flanking a sequence of stuffer DNA.
- 14. A DNA construct according to claim 12 wherein the at least two restriction sites are identical.
- 10 15. A DNA construct according to any one of the claim 12 to 13 wherein the at least one restriction site or the at least two restriction sites restriction sites are BstXI sites.
- 15 16. A DNA construct according to any one of the claims 12 to 13 wherein the restriction sites are XcmI sites.
- 17. A DNA construct according to any one of the
 20 preceding claims wherein the cloning site further
 comprises attR1 and attR2 recombination
 sequences.
- 18. A DNA construct according to any one of the
 25 preceding claims wherein the cloning site further
 comprises a ccdB nucleotide sequence.
- 19. A DNA construct according to claim 17 wherein the ccdB nucleotide sequence further comprises at
 30 least one unique restriction site.
 - 20. A DNA construct according to claim 18 wherein the at least one unique restriction site are blunt-end restriction sites.

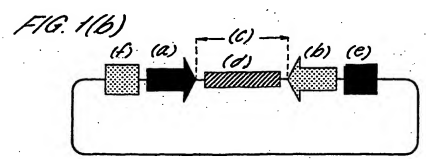
- 40 -

- 21. A DNA construct according to claim 19 wherein the blunt-end restriction sites are *Srf*I sites.
- 5 22. A DNA according to any one of the preceding claims which further comprises:
 - g) a DNA fragment inserted in the at least one cloning site.
- 10 23. A DNA construct according to any one of the preceding claims which is a plasmid or vector.

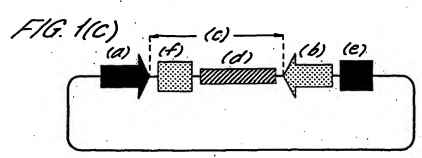
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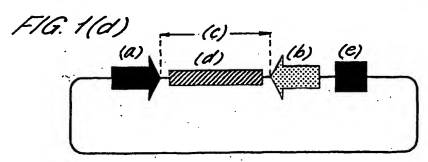
- 24. A plasmid or vector as claimed in claim 23 having the nucleotide sequence illustrated in Figure 10, Figure 11, Figure 12, Figure 13, Figure 14, or Figure 15.
- 25. Use of the DNA construct according to any one of the preceding claims for the production of double-stranded RNA for RNA inhibition.
- 26. A bacterial strain harbouring the DNA construct according to any one of the preceding claims.
- 25 27. A bacterial strain according to claim 26, wherein said bacteria strain is an *E. coli* strain.
- 28. Use of the bacterial strain according to claims
 26 or 27 for the production of double-stranded
 30 RNA for RNA inhibition.

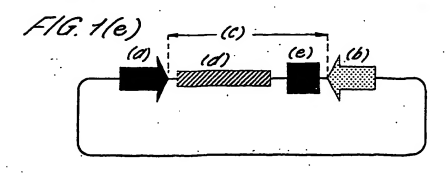


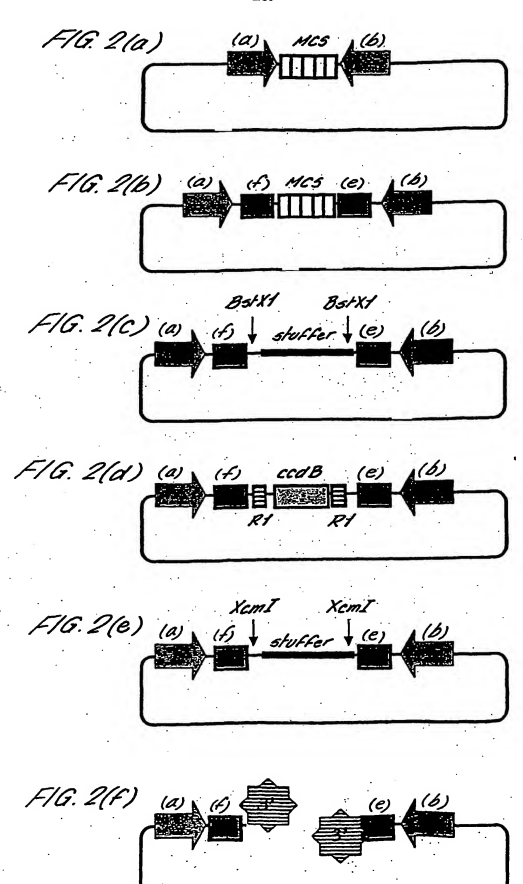


- (a): promoter 1
 (b): promoter 2
- (e): Terminator 1
 (f): Terminator 2



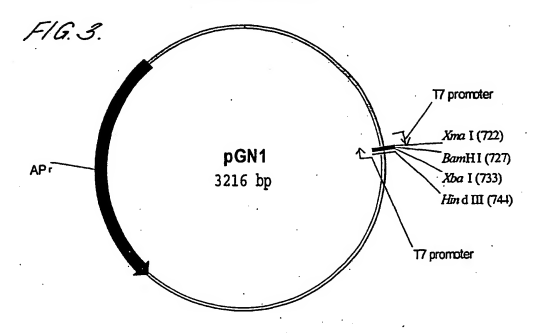


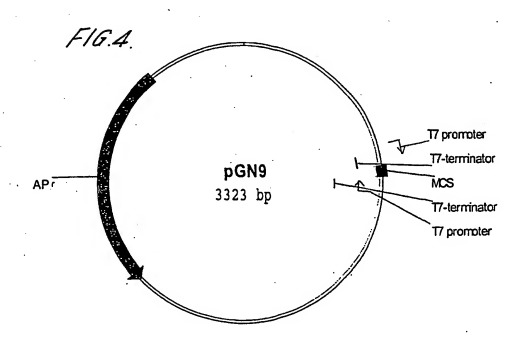




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Construction RNAi vector with T7 terminators





F16.5

T7p TTGTAATACG ACTCACTATA

631

AACATTATGC TGAGTGATAT

GGGCGAATTC GAGCTCGGTA CCCGGGGATC CTCTAGAGTC GAAAGCTTCT CGCCCTATAG TGAGTCGTAT CCCGCTTAAG CTCGAGCCAT GGGCCCCTAG GAGATCTCAG CTTTCGAAGA GCGGGATATC ACTCAGCATA 701

771 TACAGCTTGA GTATTCTATA GTGTCACCTA AATAGCTTGG CGTAATCATG GTCATAGCTG TITCCTGTGT ATGICGAACT CATAAGATAT CACAGTGGAT TTATCGAACC GCATTAGTAC CAGTATCGAC AAAGGACACA T7p

F/6.6.

T7 terminator

actag ${f cata}$ arc ${f cct}$ tagag ${f cct}$ ctaaaag ${f cct}$ ctagagat ${f ccc}$ gaaact ${f ccc}$ aaaaa ${f ccc}$

T7 terminator

ECORI COM.

KpnI EcoRI PstI 5' AATTCAAAAACCCCTCAAGACCCGTTTAGAGGCCCCAAGGGGTTATGCTAGTGAATT<u>CTGCAG</u>CGGTAC 3' oGN27

oGN28

T7 terminator

Xbal Mlul HindIII

HindIII com.

5'CTAGACGCGTAAGCTTACTAGCATAACCCCTTGGGGCCTCTAAACGGGGTCTTGAGGGGGTTTTTTG 'n oGN 29 oGN 30

TGCGCATTCGAATGATCGTATTGGGGAACCCCGGAGATTTGCCCAGGAACTCCCCAAAAACTCGA

F16.8

631

T7p

TTGTAATACG ACTCACTATA
AACATTATGC TGAGTGATAT

GGGCGAATTC AAAAAACCCC TCAAGACCCG TTTAGAGGCC CCAAGGGGTT ATGCTAGTGA ATTCTGCAGG CCCGCTTAAG TITITIGGG AGTICIGGG AAATCTCCGG GGITCCCCAA TACGATCACT TAAGACGTCC 701

T7 term

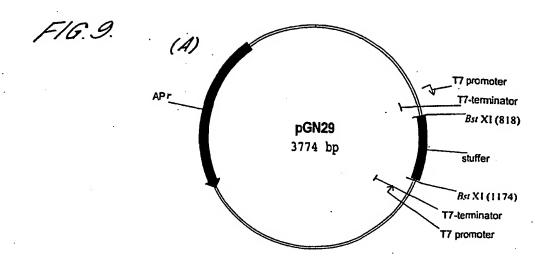
CATGGGCCCC TAGGAGATCT GCGCATTCGA ATGATCGTAT TGGGGAACCC CGGAGATTTG CCCAGAACTC GGGTTTTTTG AGCTTCTCGC CCTATAGTGA GTCGTATTAC AGCTTGAGTA TTCTATAGTG TCACCTAAAT 841

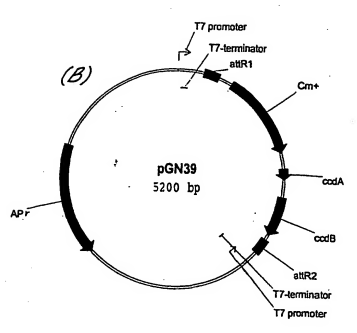
GTACCCGGGG ATCCTCTAGA CGCGTAAGCT TACTAGCATA ACCCCTTGGG GCCTCTAAAC GGGTCTTGAG

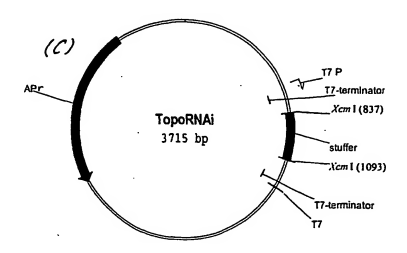
171

T7p

CCCAAAAAAC TCGAAGAGCG GGATATCACT CAGCATAATG TCGAACTCAT AAGATATCAC AGTGGATTTA







F1G. 10.

pGN9

1 gagtgcacca tatgcggtgt gaaataccgc acagatgcgt aaggagaaaa taccgcatca 61 ggcgaaattg taaacgitaa tattttgtta aaattcgcgt taaatatttg ttaaatcagc 121 teattttta accaatagge egaaategge aaaateeett ataaateaaa agaatagaee 181 gagatagggt tgagtgttgt tccagtttgg aacaagagtc cactattaaa gaacgtggac 241 tccaacgtca aagggcgaaa aaccgtctat cagggcgatg gcccactacg tgaaccatca 301 cccaaatcaa gttttttgcg gtcgaggtgc cgtaaagctc taaatcggaa ccctaaaggg 361 agecceegat ttagagettg acggggaaag ceggegaaeg tggcgagaaa ggaagggaag 421 aaagcgaaag gagcgggcgc tagggcgctg gcaagtgtag cggtcacgct gcgcgtaacc 481 accacacccg ccgcgcttaa tgcgccgcta cagggcgcgt ccattcgcca ttcaggctgc 541 gcaactgttg ggaagggcga teggtgeggg cetetteget attaegeeag etggegaaag 601 ggggatgtgc tgcaaggcga ttaagttggg taacgccagg gttttcccag tcacgacgtt 661 gtaaaacgac ggccagtgaa ttgtaatacg actcactata gggcgaattc aaaaaacccc 721 tcaagacccg tttagaggcc ccaaggggtt atgctagtga attctgcagg gtacccgggg 781 atcetetaga egegtaaget tactageata acceettggg geetetaaac gggtettgag 841 gggtttttig agettetege ectatagtga gtegtatiae agettgagta tietatagtg 901 tcacctaaat agettggcgt aatcatggtc atagetgttt cetgtgtgaa attgttatee 961 getcacaatt ccacacaaca tacgageegg aageataaag tgtaaageet ggggtgeeta 1021 atgagtgage taactcacat taattgegtt gegeteactg ecceptitee agtegggaaa 1081 cctgtcgtgc cagctgcatt aatgaatcgg ccaacgcgcg gggagaggcg gtttgcgtat 1141 tgggcgctct tccgcttcct cgctcactga ctcgctgcgc tcggtcgttc ggctgcgcg 1201 ageggtatea geteacteaa aggeggtaat aeggttatee acagaateag gggataaege 1261 aggaaagaac atgtgagcaa aaggccagca aaaggccagg aaccgtaaaa aggccgcgtt 1321 gctggcgttt ttcgataggc tccgccccc tgacgagcat cacaaaaatc gacgctcaag 1381 tcagaggtgg cgaaacccga caggactata aagataccag gcgtttcccc ctggaagctc 1441 cctcgtgcgc tctcctgttc cgaccctgcc gcttaccgga tacctgtccg cctttctccc 1501 ttcgggaage gtggcgcttt ctcatagete acgetgtagg tatetcagtt cggtgtaggt 1561 cgttcgctcc aagctgggct gtgtgcacga accecccgtt cagcccgacc gctgcgcctt 1621 atccggtaac tatcgtcttg agtccaaccc ggtaagacac gacttatcgc cactggcagc 1681 agccactggt aacaggatta gcagagcgag gtatgtaggc ggtgctacag agttcttgaa 1741 gtggtggcct aactacggct acactagaag gacagtattt ggtatctgcg ctctgctgaa 1801 gccagttacc ttcggaaaaa gagttggtag ctcttgatcc ggcaaacaaa ccaccgctgg 1861 tagcggtggt ttttttgttt gcaagcagca gattacgcgc agaaaaaaag gatctcaaga 1921 agateetttg atettteta eggggtetga egeteagtgg aacgaaaact eacgttaagg 1981 gattttggte atgagattat caaaaaggat etteacetag ateetttaa attaaaaatg 2041 aagttttaaa tcaatctaaa gtatatatga gtaaacttgg tctgacagtt accaatgctt 2101 aatcagtgag gcacctatct cagcgatctg tctatttcgt tcatccatag ttgcctgact 2161 ccccgtcgtg tagataacta cgatacggga gggcttacca tctggcccca gtgctgcaat 2221 gataccgcga gacccacgct caccggctcc agatttatca gcaataaacc agccagccgg 2281 aagggccgag cgcagaagtg gtcctgcaac titatccgcc tccatccagt ctattaattg 2341 ttgccgggaa gctagagtaa gtagttcgcc agttaatagt ttgcgcaacg ttgttggcat 2401 tgctacagge atcgtggtgt cacgetegte gtttggtatg getteattea geteeggtte 2461 ccaacgatea aggegagtta catgateece catgttgtge aaaaaagegg ttageteett 2521 cggtcctccg atcgttgtca gaagtaagtt ggccgcagtg ttatcactca tggttatggc 2581 agcactgcat aattetetta etgicatgce ateegtaaga tgettttetg tgactggtga 2641 gtactcaacc aagtcattct gagaataccg cgcccggcga ccgagttgct cttgcccggc 2701 gtcaatacgg gataatagtg tatgacatag cagaacttta aaagtgctca tcattggaaa 2761 acgttetteg gggegaaaac teteaaggat ettacegetg ttgagateca gttegatgta 2821 acccactcgt gcacccaact gatcttcagc atctttact ttcaccagcg tttctgggtg 2881 agcaaaaaca ggaaggcaaa atgccgcaaa aaagggaata agggcgacac ggaaatgttg 2941 aatactcata ctcttccttt ttcaatatta ttgaagcatt tatcagggtt attgtctcat 3001 gageggatae atatttgaat gtatttagaa aaataaacaa ataggggtte egegeacatt 3061 teccegaaaa gtgccacetg acgtetaaga aaccattatt atcatgacat taacctataa 3121 aaataggegt atcacgagge cetttegtet egegegtte ggtgatgaeg gtgaaaacet 3181 ctgacacatg cageteeegg agaeggteae agettgtetg taageggatg eegggageag 3241 acaagcccgt cagggcgcgt cagcgggtgt tggcgggtgt cggggctggc ttaactatgc 3301 ggcatcagag cagattgtac tga

FIG. 11.

PGN29

1 gagtgcacca tatgcggtgt gaaataccgc acagatgcgt aaggagaaaa taccgcatca 61 ggcgaaattg taaacgttaa tattttgtta aaattcgcgt taaatatttg ttaaatcagc 121 tcattttta accaatagge cgaaategge aaaateeett ataaateaaa agaatagace 181 gagatagggt tgagtgttgt tccagtttgg aacaagagtc cactattaaa gaacgtggac 241 tecaacetca aaggeegaaa aaceetcat caggeegate eccactace teaaceatca 301 cecaaatcaa etttttee etcaaeete cetaaageete taaateegaa cectaaagee 421 aaagcgaaag gagcggggc tagggcgctg gcaagtgtag cggtcacgct gcgcgtaacc 481 accacacccg ccgcgcttaa tgcgccgcta cagggcgcgt ccattcgcca ttcaggctgc 541 gcaactgttg ggaagggcga tcggtgcggg cctcttcgct attacgccag ctggcgaaag 601 qqqqatqtqc tqcaagqcga ttaaqttggq taacqccagq gttttcccag tcacqacqtt 661 gtaaaacgac ggccagtgaa ttgtaatacg actcactata gggcgaattc aaaaaacccc 721 tcaagacccg tttagaggcc ccaaggggtt atgctagtga attctgcagg gtacccgggg 781 atcctctaga gatccctcga cctcgagatc cattgtgctg gcgcggattc tttatcactg 841 ataagttggt ggacatatta tgtttatcag tgataaagtg tcaagcatga caaagttgca 901 gccgaataca gtgatccgtg ccggccctgg actgttgaac gaggtcggcg tagacggtct 961 gacgacacgc aaactggcgg aacggttggg ggtgcagcag ccggcgcttt actggcactt 1021 caggaacaag cgggcgctgc tcgacgcact ggccgaagcc atgctggcgg agaatcatac 1081 gcttcggtgc cgagagccga cgacgactgg cgctcatttc tgatcgggaa tcccgcagct 1141 tcaggcaggc gctgctcgcc taccgccagc acaatggatc tcgagggatc ttccatacct 1201 accagttctg cgcctgcagg tcgcggccgc gactctctag acgcgtaagc ttactagcat 1261 aacccettgg ggeetetaaa egggtettga ggggtttttt gagetteteg eectatagtg 1321 agtcgtatta cagcttgagt attctatagt gtcacctaaa tagcttggcg taatcatggt 1381 catagctgtt tcctgtgtga aattgttatc cgctcacaat tccacaacaac atacgagccg 1441 qaagcataaa gtgtaaagcc tggggtgcct aatgagtgag ctaactcaca ttaattgcgt 1501 tgcgctcact gcccgctttc cagtcgggaa acctgtcgtg ccagctgcat taatgaatcg 1561 gccaacgcgc ggggagaggc ggtttgcgta ttgggcgctc ttccgcttcc tcgctcactg 1621 actcgctgcg ctcggtcgtt cggctgcggc gagcggtatc agctcactca aaggcggtaa 1681 tacggttate cacagaatea ggggataaeg caggaaagaa catgtgagea aaaggeeage 1741 aaaaggccag gaaccgtaaa aaggccgcgt tgctggcgtt tttcgatagg ctccgcccc 1801 ctgacgagca tcacaaaaat cgacgctcaa gtcagaggtg gcgaaacccg acaggactat 1861 aaagatacca ggcgtttccc cetggaaget ceetegtgeg etetectgtt cegaccetge 1921 cgcttaccgg atacctgtcc gcctttctcc cttcgggaag cgtggcgctt tctcatagct 1981 cacgctgtag gtatctcagt tcggtgtagg tcgttcgctc caagctgggc tgtgtgcacg 2041 aacccccgt tcagcccgac cgctgcgcct tatccggtaa ctatcgtctt gagtccaacc 2101 cggtaagaca cgacttatcg ccactggcag cagccactgg taacaggatt agcagagcga 2161 ggtatgtagg cggtgctaca gagttcttga agtggtggcc taactacggc tacactagaa 2221 ggacagtatt tggtatctgc gctctgctga agccagttac cttcggaaaa agagttggta 2281 gctcttgatc cggcaaacaa accaccgctg gtagcggtgg tttttttgtt tgcaagcagc 2341 agattacgcg cagaaaaaaa ggatctcaag aagatccttt gatctttct acggggtctg 2401 acgctcagtg gaacgaaaac tcacgttaag ggattttggt catgagatta tcaaaaagga 2461 tcttcaccta gatcctttta aattaaaaat gaagttttaa atcaatctaa agtatatatg 2521 agtaaacttg gtctgacagt taccaatgct taatcagtga ggcacctatc tcagcgatct 2581 gtctatttcg ttcatccata gttgcctgac tccccgtcgt gtagataact acgatacggg 2641 agggcttacc atctggcccc agtgctgcaa tgataccgcg agacccacgc tcaccggctc 2701 cagatttate ageaataaac cageeageeg gaagggeega gegeagaagt ggteetgeaa 2761 ctttatccge ctccatccag tctattaatt gttgccggga agctagagta agtagttcgc 2821 cagttaatag tttgcgcaac gttgttggca ttgctacagg catcgtggtg tcacgctcgt 2881 cgtttggtat ggcttcattc agctccggtt cccaacgatc aaggcgagtt acatgatccc 2941 ccatgifigtg caaaaaagcg gitagcicct teggiccice gategitgic agaagtaagt 3001 tggccgcagt gttatcactc atggttatgg cagcactgca taattctctt actgtcatgc 3061 catccgtaag atgctttct gtgactggtg agtactcaac caagtcattc tgagaatacc 3121 gcgcccggcg accgagttgc tcttgcccgg cgtcaatacg ggataatagt gtatgacata 3181 gcagaacttt aaaagtgctc atcattggaa aacgttcttc ggggcgaaaa ctctcaagga 3241 tettaceget gttgagatec agttegatgt aacceaeteg tgeaeceaac tgatetteag 3301 catetttac tttcaccage gtttetgggt gagcaaaaac aggaaggeaa aatgeegeaa 3361 aaaagggaat aagggegaca eggaaatgtt gaatacteat actetteett tttcaatatt 3421 attgaagcat ttatcagggt tattgtctca tgagcggata catatttgaa tgtatttaga 3481 aaaataaaca aataggggtt ccgcgcacat ttccccgaaa agtgccacct gacgtctaag 3541 aaaccattat tatcatgaca ttaacctata aaaataggcg tatcacgagg ccctttcgtc 3601 tcgcgcgttt cggtgatgac ggtgaaaacc tctgacacat gcagctcccg gagacggtca 3661 cagettgtet gtaageggat geegggagea gacaageeeg teagggegeg teagegggtg 3721 ttggcgggtg tcggggctgg cttaactatg cggcatcaga gcagattgta ctga

F16.12.

pGN39

TAATACGACT CACTATAGGG CGAATTCAAA AAACCCCTCA AGACCCGTTT AGAGGCCCCA AGGGGTTATG CTAGTGAATT CTGCAGCGGT ACCCGGGGAT CCTCTAGAGA TCCCTCGACC TCGAGATCCA TTGTGCTGGA AAGATCACAA GTTTGTACAA AAAAGCTGAA CGAGAAACGT AAAATGATAT AAATATCAAT ATATTAAATT AGATTTTGCA TAAAAAACAG ACTACATAAT ACTGTAAAAC ACAACATATC CAGTCACTAT GGCGGCCGCA TTAGGCACCC CAGGCTTTAC ACTITATGCT TCCGGCTCGT ATAATGTGTG GATTTTGAGT TAGGATCCGG CGAGATTTTC AGGAGCTAAG GAAGCTAAAA TGGAGAAAAA AATCACTGGA TATACCACCG TTGATATATC CCAATGGCAT CGTAAAGAAC ATTTTGAGGC ATTTCAGTCA GTTGCTCAAT GTACCTATAA CCAGACCGTT CAGCTGGATA TTACGGCCTT TTTAAAGACC GTAAAGAAAA ATAAGCACAA GTTTTATCCG GCCTTTATTC ACATTCTTGC CCGCCTGATG AATGCTCATC CGGAATTCCG TATGGCAATG AAAGACGGTG AGCTGGTGAT ATGGGATAGT GTTCACCCTT GTTACACCGT TTTCCATGAG CAAACTGAAA CGTTTTCATC GCTCTGGAGT GAATACCACG ACGATTTCCG GCAGTTTCTA CACATATATT CGCAAGATGT GGCGTGTTAC GGTGAAAACC TGGCCTATTT CCCTAAAGGG TTTATTGAGA ATATGTTTTT CGTCTCAGCC AATCCCTGGG TGAGTTTCAC CAGTTTTGAT TTAAACGTGG CCAATATGGA CAACTTCTTC GCCCCCGTTT TCACCATGGG CAAATATTAT ACGCAAGGCG ACAAGGTGCT GATGCCGCTG GCGATTCAGG TTCATCATGC CGTCTGTGAT GGCTTCCATG TCGGCAGAAT GCTTAATGAA TTACAACAGT ACTGCGATGA GTGGCAGGGC GGGGCGTAAA GATCTGGATC CGGCTTACTA AAAGCCAGAT AACAGTATGC GTATTTGCGC GCTGATTTTT GCGGTATAAG AATATATACT GATATGTATA CCCGAAGTAT GTCAAAAAGA GGTGTGCTAT GAAGCAGCGT ATTACAGTGA CAGTTGACAG CGACAGCTAT CAGTTGCTCA AGGCATATAT GATGTCAATA TCTCCGGTCT GGTAAGCACA ACCATGCAGA ATGAAGCCCG TCGTCTGCGT GCCGAACGCT GGAAAGCGGA AAATCAGGAA GGGATGGCTG AGGTCGCCCG GTTTATTGAA ATGAACGGCT CTTTTGCTGA CGAGAACAGG GACTGGTGAA ATGCAGTTTA AGGTTTACAC CTATAAAAGA GAGAGCCGTT ATCGTCTGTT TGTGGATGTA CAGAGTGATA TTATTGACAC GCCCGGGCGA CGGATGGTGA TCCCCCTGGC CAGTGCACGT CTGCTGTCAG ATAAAGTCTC CCGTGAACTT TACCCGGTGG TGCATATCGG GGATGAAAGC TGGCGCATGA TGACCACCGA TATGGCCAGT GTGCCGGTCT CCGTTATCGG GGAAGAAGTG GCTGATCTCA GCCACCGCGA AAATGACATC AAAAACGCCA TTAACCTGAT GTTCTGGGGA ATATAAATGT CAGGCTCCCT TATACACAGC CAGTCTGCAG GTCGACCATA GTGACTGGAT ATGTTGTGTT TTACAGTATT ATGTAGTCTG TTTTTTATGC AAAATCTAAT TTAATATATT GATATITATA TCATTITACG TITCTCGTTC AGCTITCTTG TACAAAGTGG TGATCITTCC AGCACAATGG ATCTCGAGGG ATCTTCCATA CCTACCAGTT CTGCGCCTGC AGGTCGCGGC CGCGACTCTA GACGCGTAAG CTTACTAGCA TAACCCCTTG GGGCCTCTAA ACGGGTCTTG AGGGGTTTTT TGAGCTTCTC GCCCTATAGT GAGTCGTATT ACAGCTTGAG TATTCTATAG TGTCACCTAA ATAGCTTGGC GTAATCATGG TCATAGCTGT TTCCTGTGTG AAATTGTTAT CCGCTCACAA TTCCACACAA CATACGAGCC GGAAGCATAA AGTGTAAAGC

FIG. 12 (CONTINUED 1)

CTGGGGTGCC TAATGAGTGA GCTAACTCAC ATTAATTGCG TTGCGCTCAC TGCCCGCTTT CCAGTCGGGA AACCTGTCGT GCCAGCTGCA TTAATGAATC GGCCAACGCG CGGGGAGAGG CGGTTTGCGT ATTGGGCGCT CTTCCGCTTC CTCGCTCACT GACTCGCTGC GCTCGGTCGT TCGGCTGCGG CGAGCGGTAT CAGCTCACTC AAAGGCGGTA ATACGGTTAT CCACAGAATC AGGGGATAAC GCAGGAAAGA ACATGTGAGC AAAAGGCCAG CAAAAGGCCA GGAACCGTAA AAAGGCCGCG TTGCTGGCGT TTTTCGATAG GCTCCGCCCC CCTGACGAGC ATCACAAAAA TCGACGCTCA AGTCAGAGGT GGCGAAACCC GACAGGACTA TAAAGATACC AGGCGTTTCC CCCTGGAAGC TCCCTCGTGC GCTCTCCTGT TCCGACCCTG CCGCTTACCG GATACCTGTC CGCCTTTCTC CCTTCGGGAA GCGTGGCGCT TTCTCATAGC TCACGCTGTA GGTATCTCAG TTCGGTGTAG GTCGTTCGCT CCAAGCTGGG CTGTGTGCAC GAACCCCCCG TTCAGCCCGA CCGCTGCGCC TTATCCGGTA ACTATCGTCT TGAGTCCAAC CCGGTAAGAC ACGACTTATC GCCACTGGCA GCAGCCACTG GTAACAGGAT TAGCAGAGCG AGGTATGTAG GCGGTGCTAC AGAGTTCTTG AAGTGGTGGC CTAACTACGG CTACACTAGA AGGACAGTAT TTGGTATCTG CGCTCTGCTG AAGCCAGTTA CCTTCGGAAA AAGAGTTGGT AGCTCTTGAT CCGGCAAACA AACCACCGCT GGTAGCGGTG GTTTTTTGT TTGCAAGCAG CAGATTACGC GCAGAAAAA AGGATCTCAA GAAGATCCTT TGATCTTTTC TACGGGGTCT GACGCTCAGT GGAACGAAAA CTCACGTTAA GGGATTTTGG TCATGAGATT ATCAAAAAGG ATCTTCACCT AGATCCTTTT AAATTAAAAA TGAAGTTTTA AATCAATCTA AAGTATATAT GAGTAAACTT GGTCTGACAG TTACCAATGC TTAATCAGTG AGGCACCTAT CTCAGCGATC TGTCTATTTC GTTCATCCAT AGTTGCCTGA CTCCCCGTCG TGTAGATAAC TACGATACGG GAGGGCTTAC CATCTGGCCC CAGTGCTGCA ATGATACCGC GAGACCCACG CTCACCGGCT CCAGATTTAT CAGCAATAAA CCAGCCAGCC GGAAGGGCCG AGCGCAGAAG TGGTCCTGCA ACTITATCCG CCTCCATCCA GTCTATTAAT TGTTGCCGGG AAGCTAGAGT AAGTAGTTCG CCAGTTAATA GTTTGCGCAA CGTTGTTGGC ATTGCTACAG GCATCGTGGT GTCACGCTCG TCGTTTGGTA TGGCTTCATT CAGCTCCGGT TCCCAACGAT CAAGGCGAGT TACATGATCC CCCATGTTGT GCAAAAAAGC GGTTAGCTCC TTCGGTCCTC CGATCGTTGT CAGAAGTAAG TTGGCCGCAG TGTTATCACT CATGGTTATG GCAGCACTGC ATAATTCTCT TACTGTCATG CCATCCGTAA GATGCTTTTC TGTGACTGGT GAGTACTCAA CCAAGTCATT CTGAGAATAC CGCGCCCGGC GACCGAGTTG CTCTTGCCCG GCGTCAATAC GGGATAATAG TGTATGACAT AGCAGAACTT TAAAAGTGCT CATCATTGGA AAACGTTCTT CGGGGCGAAA ACTCTCAAGG ATCTTACCGC TGTTGAGATC CAGTTCGATG TAACCCACTC GTGCACCCAA CTGATCTTCA GCATCTTTTA CTTTCACCAG CGTTTCTGGG TGAGCAAAAA CAGGAAGGCA AAATGCCGCA AAAAAGGGAA TAAGGGCGAC ACGGAAATGT TGAATACTCA TACTCTTCCT TTTTCAATAT TATTGAAGCA TTTATCAGGG TTATTGTCTC ATGAGCGGAT ACATATTTGA ATGTATTTAG AAAAATAAAC AAATAGGGGT TCCGCGCACA TTTCCCCGAA AAGTGCCACC TGACGTCTAA GAAACCATTA TTATCATGAC ATTAACCTAT AAAAATAGGC GTATCACGAG GCCCTTTCGT CTCGCGCGTT TCGGTGATGA CGGTGAAAAC CTCTGACACA TGCAGCTCCC GGAGACGGTC ACAGCTTGTC TGTAAGCGGA TGCCGGGAGC AGACAAGCCC GTCAGGGCGC GTCAGCGGGT GTTGGCGGGT GTCGGGGCTG GCTTAACTAT GCGGCATCAG

FIG. 12 (CONTINUED 2)

AGCAGATTGT ACTGAGAGTG CACCATATGC GGTGTGAAAT ACCGCACAGA
TGCGTAAGGA GAAAATACCG CATCAGGCGA AATTGTAAAC GTTAATATTT
TGTTAAAATT CGCGTTAAAT ATTTGTTAAA TCAGCTCATT TTTTAACCAA
TAGGCCGAAA TCGGCAAAAT CCCTTATAAA TCAAAAGAAT AGACCGAGAT
AGGGTTGAGT GTTGTTCCAG TTTGGAACAA GAGTCCACTA TTAAAAGAACG
TGGACTCCAA CGTCAAAGGG CGAAAAACCG TCTATCAGGG CGATGGCCCA
CTACGTGAAC CATCACCCAA ATCAAGTTTT TTGCGGTCGA GGTGCCGTAA
AGCTCTAAAT CGGAACCCTA AAGGGAGCCC CCGATTTAGA GCTTGACGGG
GAAAGCCGGC GAACGTGGCG AGAAAGGAAG GGAAGAAAGC GAAAGGAGCG
GGCGCTAGGG CGCTGGCAAG TGTAGCGGTC ACCCTGCGCG TAACCACCAC
ACCCGCCGCG CTTAATGCGC CGCTACAGGG CGCGTCCATT CGCCATTCAG
GCTGCGCAAC TGTTGGGAAG GGCGATCGGT GCGGGCCTCT TCGCTATTAC
GCCAGCTGGC GAAAGGGGGA TGTGCTGCAA GGCGATTAAG TTGGGTAACG
CCAGGGTTTT CCCCAGTCACG ACGTTGTAAA ACGACGCCA GTGAATTG

FIG. 13.

TopoRNAi

```
1 gagtgcacca tatgcggtgt gaaataccgc acagatgcgt aaggagaaaa taccgcatca
   61 ggcgaaattg taaacgttaa tattttgtta aaattcgcgt taaatatttg ttaaatcagc
  121 tcattttta accaataggc cgaaatcggc aaaatccctt ataaatcaaa agaatagacc
 181 gagatagggt tgagtgttgt tccagtttgg aacaagagtc cactattaaa gaacgtggac 241 tccaacgtca aagggcgaaa aaccgtctat cagggcgatg gcccactacg tgaaccatca 301 cccaaatcaa gttttttgcg gtcgaggtgc cgtaaagctc taaatcggaa ccctaaaggg
 541 gcaactgttg ggaagggcga tcggtgcggg cctcttcgct attacgccag ctggcgaaag 601 ggggatgtgc tgcaaggcga ttaagttggg taacgccagg gttttcccag tcacgacgtt 661 gtaaaacgac ggccagtgaa ttgtaatacg actcactata gggcgaattc aaaaaacccc
 721 tcaagacccg titagaggcc ccaaggggtt atgctagtga attctgcagg gtacccgggg
 781 atcctctaga gatccctcga cctcgagatc cattgtggtg gaattctacc aaggctagca 841 tgggcagccg aatacagtga tccgtgccgg ccctggactg ttgaacgagg tcggcgtaga 901 cggtctgacg acacgcaaac tggcggaacg gttgggggtg cagcagccgg cgctttactg
  961 gcacttcagg aacaageggg cgctgctcga cgcactggcc gaagccatgc tggcggagaa
1021 tcatacgctt cggtgccgag agccgacgac gactggcgct catttctgat cgggaatccc 1081 gcagccatgc tagccttggt agaattccac cacaatggat ctcgagggat cttccatacc
1141 taccagttet gegeetgeag gtegeggeeg egacteteta gaegegtaag ettactagea
1201 taaccccttg gggcctctaa acgggtcttg aggggttttt tgagcttctc gccctatagt 1261 gagtcgtatt acagcttgag tattctatag tgtcacctaa atagcttggc gtaatcatgg 1321 tcatagctgt ttcctgtgtg aaattgttat ccgctcacaa ttccacacaa catacgagcc
1381 ggaagcataa agtgtaaagc ctggggtgcc taatgagtga gctaactcac attaattgcg
1441 tigegeteac tgeeegetit ecagiegga aacetgiegt gecagetgea ttaatgaate
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pGN49A F/G. 14.

TGTAATACGA CTCACTATAG GGCGAATTCA AAAAACCCCT CAAGACCCGT TTAGAGGCCC CAAGGGGTTA TGCTAGTGAA TTCTGCAGCG GTACCCGGGG ATCCTCTAGA GATCCCTCGA CCTCGAGATC CATTGTGCTG GAAAGGATCT GGATCCGGCT TACTAAAAGC CAGATAACAG TATGCGTATT TGCGCGCTGA TTTTTGCGGT ATAAGAATAT ATACTGATAT GTATACCCGA AGTATGTCAA AAAGAGGTGT GCTATGAAGC AGCGTATTAC AGTGACAGTT GACAGCGACA GCTATCAGTT GCTCAAGGCA TATATGATGT CAATATCTCC GGTCTGGTAA GCACAACCAT GCAGAATGAA GCCCGTCGTC TGCGTGCCGA ACGCTGGAAA GCGGAAAATC AGGAAGGGAT GGCTGAGGTC GCCCGGTTTA TTGAAATGAA CGGCTCTTTT GCTGACGAGA ACAGGGACTG GTGAAATGCA GTTTAAGGTT TACACCTATA AAAGAGAGAG CCGTTATCGT CTGTTTGTGG ATGTACAGAG TGATATTATT GACACGCCG GGCGACGGAT GGTGATCCCC CTGGCCAGTG CACGTCTCTT AAGCGATAAA GTCTCCCGTG AACTTTACCC GGTGGTGCAT ATCGGGGATG AAAGCTGGCG CATGATGACC ACCGATATGG CCAGTGTGCC GGTCTCCGTT ATCGGGGAAG AAGTGGCTGA TCTCAGCCAC CGCGAAAATG ACATCAAAAA CGCCATTAAC CTGATGTTCT GGGGAATATA AATGTCAGGC TCCCTTATAC ACAGCCTTTC CAGCACAATG GATCTCGAGG GATCTTCCAT ACCTACCAGT TCTGCGCCTG CAGGTCGCGG CCGCGACTCT AGACGCGTAA GCTTACTAGC ATAACCCCTT GGGGCCTCTA AACGGGTCTT GAGGGGTTTT TTGAGCTTCT CGCCCTATAG TGAGTCGTAT TACAGCTTGA GTATTCTATA GTGTCACCTA AATAGCTTGG CGTAATCATG GTCATAGCTG TTTCCTGTGT GAAATTGTTA TCCGCTCACA ATTCCACACA ACATACGAGC CGGAAGCATA AAGTGTAAAG CCTGGGGTGC CTAATGAGTG AGCTAACTCA CATTAATTGC GTTGCGCTCA CTGCCCGCTT TCCAGTCGGG AAACCTGTCG TGCCAGCTGC ATTAATGAAT CGGCCAACGC GCGGGGAGAG GCGGTTTGCG TATTGGGCGC TCTTCCGCTT CCTCGCTCAC TGACTCGCTG CGCTCGGTCG TTCGGCTGCG GCGAGCGGTA TCAGCTCACT CAAAGGCGGT AATACGGTTA TCCACAGAAT CAGGGGATAA CGCAGGAAAG AACATGTGAG CAAAAGGCCA GCAAAAGGCC AGGAACCGTA AAAAGGCCGC GTTGCTGGCG TTTTTCGATA GGCTCCGCCC CCCTGACGAG CATCACAAAA ATCGACGCTC AAGTCAGAGG TGGCGAAACC CGACAGGACT ATAAAGATAC CAGGCGTTTC CCCCTGGAAG CTCCCTCGTG CGCTCTCCTG TTCCGACCCT GCCGCTTACC GGATACCTGT CCGCCTTTCT CCCTTCGGGA AGCGTGGCGC TTTCTCATAG CTCACGCTGT AGGTATCTCA GTTCGGTGTA GGTCGTTCGC TCCAAGCTGG GCTGTGTGCA CGAACCCCCC GTTCAGCCCG ACCGCTGCGC CTTATCCGGT AACTATCGTC TTGAGTCCAA CCCGGTAAGA CACGACTTAT CGCCACTGGC AGCAGCCACT GGTAACAGGA TTAGCAGAGC GAGGTATGTA GGCGGTGCTA CAGAGTTCTT GAAGTGGTGG CCTAACTACG GCTACACTAG AAGGACAGTA TTTGGTATCT GCGCTCTGCT GAAGCCAGTT ACCTTCGGAA AAAGAGTTGG TAGCTCTTGA TCCGGCAAAC AAACCACCGC TGGTAGCGGT GGTTTTTTTG TTTGCAAGCA GCAGATTACG CGCAGAAAAA AAGGATCTCA AGAAGATCCT TTGATCTTTT CTACGGGGTC TGACGCTCAG TGGAACGAAA ACTCACGTTA AGGGATTTTG GTCATGAGAT TATCAAAAG GATCTTCACC TAGATCCTTT TAAATTAAAA ATGAAGTTTT

FIG. 14 (CONTINUED)

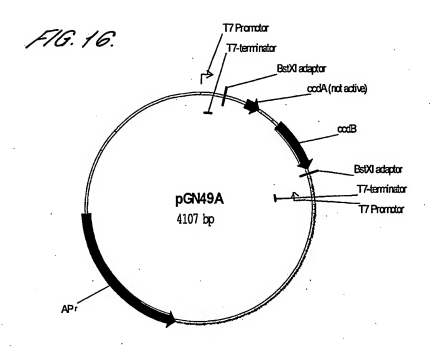
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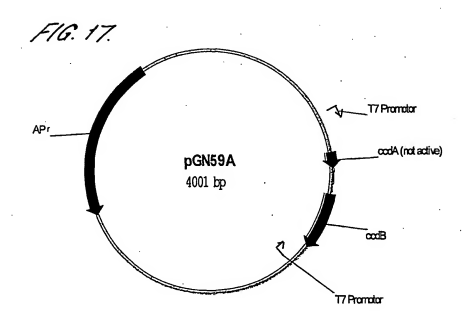
PGN59A FIG. 15.

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FIG. 15 (CONTINUED)

CATAGCTCAC GCTGTAGGTA TCTCAGTTCG GTGTAGGTCG TTCGCTCCAA GCTGGGCTGT GTGCACGAAC CCCCCGTTCA GCCCGACCGC TGCGCCTTAT CCGGTAACTA TCGTCTTGAG TCCAACCCGG TAAGACACGA CTTATCGCCA CTGGCAGCAG CCACTGGTAA CAGGATTAGC AGAGCGAGGT ATGTAGGCGG TGCTACAGAG TTCTTGAAGT GGTGGCCTAA CTACGGCTAC ACTAGAAGGA CAGTATTTGG TATCTGCGCT CTGCTGAAGC CAGTTACCTT CGGAAAAAGA GTTGGTAGCT CTTGATCCGG CAAACAAACC ACCGCTGGTA GCGGTGGTTT TTTTGTTTGC AAGCAGCAGA TTACGCGCAG AAAAAAAGGA TCTCAAGAAG ATCCTTTGAT CTTTTCTACG GGGTCTGACG CTCAGTGGAA CGAAAACTCA CGTTAAGGGA TTTTGGTCAT GAGATTATCA AAAAGGATCT TCACCTAGAT CCTTTTAAAT TAAAAATGAA GTTTTAAATC AATCTAAAGT ATATATGAGT AAACTTGGTC TGACAGTTAC CAATGCTTAA TCAGTGAGGC ACCTATCTCA GCGATCTGTC TATTTCGTTC ATCCATAGTT GCCTGACTCC CCGTCGTGTA GATAACTACG ATACGGGAGG GCTTACCATC TGGCCCCAGT GCTGCAATGA TACCGCGAGA CCCACGCTCA CCGGCTCCAG ATTTATCAGC AATAAACCAG CCAGCCGGAA GGGCCGAGCG CAGAAGTGGT CCTGCAACTT TATCCGCCTC CATCCAGTCT ATTAATTGTT GCCGGGAAGC TAGAGTAAGT AGTTCGCCAG TTAATAGTTT GCGCAACGTT GTTGGCATTG CTACAGGCAT CGTGGTGTCA CGCTCGTCGT TTGGTATGGC TTCATTCAGC TCCGGTTCCC AACGATCAAG GCGAGTTACA TGATCCCCCA TGTTGTGCAA AAAAGCGGTT AGCTCCTTCG GTCCTCCGAT CGTTGTCAGA AGTAAGTTGG CCGCAGTGTT ATCACTCATG GTTATGGCAG CACTGCATAA TTCTCTTACT GTCATGCCAT CCGTAAGATG CTTTTCTGTG ACTGGTGAGT ACTCAACCAA GTCATTCTGA GAATACCGCG CCCGGCGACC GAGTTGCTCT TGCCCGGCGT CAATACGGGA TAATAGTGTA TGACATAGCA GAACTTTAAA AGTGCTCATC ATTGGAAAAC GTTCTTCGGG GCGAAAACTC TCAAGGATCT TACCGCTGTT GAGATCCAGT TCGATGTAAC CCACTCGTGC ACCCAACTGA TCTTCAGCAT CTTTTACTTT CACCAGCGTT TCTGGGTGAG CAAAAACAGG AAGGCAAAAA GCCGCAAAAA AGGGAATAAG GGCGACACGG AAATGTTGAA TACTCATACT CTTCCTTTTT CAATATTATT GAAGCATTTA TCAGGGTTAT TGTCTCATGA GCGGATACAT ATTTGAATGT ATTTAGAAAA ATAAACAAAT AGGGGTTCCG CGCACATTTC CCCGAAAAGT GCCACCTGAC GTCTAAGAAA CCATTATTAT CATGACATTA ACCTATAAAA ATAGGCGTAT CACGAGGCCC TTTCGTCTCG CGCGTTTCGG TGATGACGGT GAAAACCTCT GACACATGCA GCTCCCGGAG ACGGTCACAG CTTGTCTGTA AGCGGATGCC GGGAGCAGAC AAGCCCGTCA GGGCGCGTCA GCGGGTGTTG GCGGGTGTCG GGGCTGGCTT AACTATGCGG CATCAGAGCA GATTGTACTG A





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INTERNATIONAL SEARCH REPORT

Hional Application No PCI/IB 01/01068

A. CLASSIFICATION OF SUBJECT MATTER IPC 7 C12N15/10 C12N15/63 C12N15/70 C12N1/21 According to International Patent Classification (IPC) or to both national classification and IPC Minimum documentation searched (classification system followed by classification symbols) IPC 7 C12N Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) EPO-Internal, WPI Data, BIOSIS, CHEM ABS Data, MEDLINE C. DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages Relevant to dalm No. Category ° X FR 2 782 325 A (PROTEUS) 1-24,26, 18 February 2000 (2000-02-18) page 7, line 20 -page 8, line 8 page 11, line 11 - line 36 page 23, line 31 -page 24, line 9 1 - 28WO 00 01846 A (DEVGEN N.V.) A 13 January 2000 (2000-01-13) cited in the application page 8, line 9 -page 10, line 22 page 15, line 9 - line 33 page 21, line 21 -page 22, line 29 Patent family members are listed in annex. Further documents are listed in the continuation of box C. Special categories of cited documents: 'T' later document published after the International filing date or priority date and not in conflict with the application but clied to understand the principle or theory underlying the "A" document defining the general state of the art which is not considered to be of particular relevance 'E' earlier document but published on or after the international "X" document of particular relevance; the claimed Invention cannot be considered novel or cannot be considered to 'L' document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) involve an inventive step when the document is taken alone "Y" document of particular relevance; the ctaimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such docu-ments, such combination being obvious to a person skilled 'O' document referring to an oral disclosure, use, exhibition or other means document published prior to the International filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the International search report 20 September 2001 27/09/2001 Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Montero Lopez, B Fax: (+31-70) 340-3016

INTERNATIONAL SEARCH REPORT

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Continua	tion) DOCUMENTS CONSIDERED TO BE RELEVANT	· · · · · · · · · · · · · · · · · · ·		
ategory °	Citation of document, with indication, where appropriate, of the relevant passages		Relevant to claim	No.
·,Х	WO 01 34815 A (CAMBRIA BIOSCIENCES, LLC) 17 May 2001 (2001-05-17) page 5, last paragraph -page 6, paragraph 4 page 20, paragraph 2 page 24, last paragraph; example 1 page 13, last paragraph -page 15, paragraph 2		1-10, 22,23	12, ,25
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INTERNATIONAL SEARCH REPORT

information on patent family members

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